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MAUDE Adverse Event Report: EPIC EPIC EHR SOFTWARE

6510(k)⁷|DeNovo⁸|Registration & Listing⁹

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CDRH Supersearch CERTI

Events¹⁰

CFR Title 21¹⁶ Radiation-Emitting Products 17 X-Ray Assembler 18 Medsun Reports 19 CLIA 20 TPLC 21

EPIC EPIC EHR SOFTWARE

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Event Date 06/09/2017 Event Type Injury Event Description

Pt presented to emergency ward with delirium and leukocytosis. Extensive neurological eval was undertaken. It was negative. The chest radiograph with reported late, and the report came back silently into the radiograph silo, as pneumonia, but no one saw the report. There was not any notice or warning that a new report had been generated. In addition, the ehr device uses elaborate system to generate a complete history and physical exam report. There was no way that the lungs were "clear" as stated by the doctors (er and attending) using the canned language of the ehr macro/template when the pt had bilateral pneumonia. The treatment with antibiotics was delayed by 20 hours. This case raises important issues that have been wrought by ehr devices that have not had any vetting for safety, unstability, and efficacy, and remain free of after market surveillance. Reports of all types get deposited into their respective silos silently, and no one knows the results for hours whether they are good or bad. Care is delayed with frequent life threatening consequences. The ehr device enables elaborate reports of examinations, that often are not done, with one swift click of the mouse. Basically, these are fake exams and histories.

Search Alerts/Recalls²²

New Search | Submit an Adverse Event Report²³

Brand NameEPIC EHR
Type of DeviceSOFTWARE
Manufacturer (Section D)EPIC

Verona WI 53593

MDR Report Key6636342 Report NumberMW5070360

Device Sequence Number1

Product CodeJQP24

Report Source Voluntary

Reporter Occupation Physician

Report Date06/10/2017

1 Device Was Involved in the Event

1 Patient Was Involved in the Event

Date FDA Received06/10/2017

Is This An Adverse Event Report?Yes

Is This A Product Problem Report?Yes

Device OperatorHealth Professional

Was Device Available For Evaluation?Yes

Is The Reporter A Health Professional?Yes

Was the Report Sent to FDA?

Event LocationNo Information

Was Device Evaluated By Manufacturer?

Is The Device Single Use?

Is this a Reprocessed and Reused Single-Use Device?No

Type of Device Usage

Patient TREATMENT DATA

Date Received: 06/10/2017 Patient Sequence Number: 1

Links on this page:

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- 20. /scripts/cdrh/cfdocs/cfClia/Search.cfm
- 21. /scripts/cdrh/cfdocs/cfTPLC/tplc.cfm
- 22. https://www.fda.gov/MedicalDevices/Safety/ListofRecalls/default.htm
- 23. https://www.accessdata.fda.gov/scripts/medwatch/
- 24. ../cfPCD/classification.cfm?start_search=&ProductCode=JQP

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